

Amendments to the specification:

- In the specification after page 24, replace the Sequence Listing with the substitute Sequence Listing on pages 1-8 enclosed herewith.

In the following, added portions are indicated by **underlining and boldface**, to avoid confusion with portions underlined in the original text.

- On page 14, please amend the paragraph starting on line 19 as follows:

In this example, an oligonucleotide tag repertoire is produced such that each oligonucleotide tag consists of eight words of four nucleotides. The procedure outlined in Figure 2 is followed. A vector, corresponding to vector (200), is constructed by first inserting the following oligonucleotide (SEQ ID NO: 4 and 27, respectively) into a Bam HI and Eco RI digested pUC19:

PacI		Bse RI	Bsp 120		Bbs I	Eco RI	Bam HI
↓		↓	↓		↓	↓	↓
aattg <u>tt</u> aatt <u>aagg</u> atgagctcactcctc <u>gggcccgc</u> cataagtcttcga <u>attcg</u>							
caattaattcctactcgagtg <u>aggagcccgggc</u> gtattc <u>agaagctt</u> aagcctag							

Formula II

Separately, the oligonucleotide of Formula I and forward and reverse primers (SEQ ID NO: 2) and SEQ ID NO: 3) are synthesized using a conventional DNA synthesizer, e.g. PE Applied Biosystems (Foster City, CA) model 392. The oligonucleotide of Formula I is a mixture containing a repertoire of 64 two-word oligonucleotide tag precursors. The four-nucleotide words of Table I are employed. After amplification by PCR[,] the amplification product is digested with Bbs I to give the following two products (SEQ ID NO: 28 and 29, **respectively**):

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... gaagacga          word-word-gg ...
... cttctgct-word      word-cc ...

```

The products are re-ligated, amplified by PCR, and digested with Bbv I to give the following two products (SEQ ID NO: 30 and 31, respectively):

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... gaagacga-word      word-gg ...
... cttctgct-word-word  cc ...

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The products are again re-ligated and amplified by PCR. By this sequence of cleavages and relations, any words consisting of failure sequences are selected against by the ligation event, i.e. words with failure sequences will not religate in the mixture, and thus, will not be amplified. The final product is digested with Pst I and Hind III and inserted into a Pst I/Hind III-digested pUC19 To give the following construct (SEQ ID NO: 5):

Pst I	Bse RI	Bbs I		Bsp 120		Hind III
↓	↓	↓		↓		↓
...cgacctgcagaggagatgaagacga-wordword-gggcccaatgctgcaagcttggcg...						
...gctggacgtctcctctacttctgct-wordword-cccgggttacgacgttcgaaccgc...						
					↑	
					Bbv I	

where Pst I, Bse RI, Bbs I, Bsp 120, and Bbv I, correspond to r_4 , r_5 , r_6 , r_7 , and r_8 of Figure 2, respectively. After amplification in a suitable host, the plasmid is isolated and cleaved with Pst I and Bbs I to give an opened vector with the following upstream and downstream (SEQ ID NO: 6 and 32, respectively) ends:

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...cgacctgca      wordword-gggcccaatgctgcaagcttggcg...
...gctgg          word-cccgggttacgacgttcgaaccgc...

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Separately, a portion of the amplified oligonucleotide of Formula I is digested with Pst I and Bbv I to give the following fragment (SEQ ID NO: 7 and 38, respectively):

gaggagatgaagacga-word
acgtctcctctacttctgct-wordword

This fragment is inserted into the above vector opened by digestion with Bbs I and Pst I to give the following construct (SEQ ID NO: 8):

...gcagaggagatgaagacga-wordwordword-gggcccaatgctgcaagcttggcg...
...cgtctcctctacttctgct-wordwordword-cccgggttacgacgttcgaaccgc...

which contains an oligonucleotide tag precursor of three words. The steps of cleaving, inserting, and amplification are repeated until a construct containing eight words is obtained. Preferably, at each step, reactants, e.g. vectors and/or inserts, are provided in amounts that are at least ten times the complexity of the reactant. When synthesis is complete, the eight-word construct is cleaved with Bse RI and Bsp 120 and the following fragment containing the oligonucleotide tag repertoire is isolated (complement is SEQ ID NO:33):

(word)₈g
ct (word)₈cccgg

The isolated fragment is then inserted into the Bse RI/Bsp 120 vector of Formula II, which vector is used to transform a suitable host. The construct is ready for inserting polynucleotides, such as cDNAs, into the Eco RI restriction site to form tag-polynucleotide conjugates in accordance with the method of Brenner et al., International patent application pct/us96/09513.

- Please amend the paragraph starting at page 17, line 30 as follows:

After cloning, the population of vectors is divided into two parts, after which the vectors in one part are cleaved with Pst I and Bsg I to give the following fragment mixture (SEQ ID NO:11 and 34, respectively):

gttatcggaggagatgaagacgg [word] [word] gg
acgtcaatagcctcctctacttctgcc [word] [word]

which is isolated. The vectors in the other part are cleaved with Pst I and Bse RI and the linearized word-containing vectors are isolated. The word-containing fragments are ligated into the linearized vectors to form the following construct (SEQ ID NO: 12):

...ctgcagttatcggaggagatgaagacgg [word] [word] gg [word] [word] -
...gacgtcaatagcctcctctacttctgcc [word] [word] cc [word] [word] -

-gggcccataatatccgtctgcacaagcttaeggcg...
-cccggtatataggcagacgtgttcgaaccgc...

After cloning, the construct is again divided into two parts and the steps are repeated to give the final 8-word repertoire having the form (SEQ ID NO: 35):

.. gaagacgg ([word] [word] gg) ₄gcc ...
.. cttctgcc ([word] [word] cc) ₄cgg ...

This may then be cleaved with Bse RI and Bsg I and re-cloned into a vector similar to that of Formula II for attachment to polynucleotides.

Please amend the paragraph starting at page 18, line 36 as follows:

pUC19 was digested to completion with Sap I and Eco RI using the manufacturer's protocol and the large fragment was isolated. All restriction endonucleases unless otherwise noted were

purchased from New England Biolabs (Beverly, MA). The small Sap I-Eco RI fragment was removed to eliminate the β -gal promoter sequence, which was found to skew the representation of some combinations of words in the final library. The following adaptor (SEQ ID NO: 13 and 36, respectively) was ligated to the isolated large fragment in a conventional ligation reaction to give plasmid pUCSE as a ligation product.

```

Eco RI      Pst I      Eco RV Hind III
  ↓          ↓          ↓          ↓
aattctagactgcagttgatatcttaagctt
      gatctgacgtcaactatagaattcgaacga

```

A bacterial host was transformed by the ligation product using electroporation, after which the transformed bacteria were plated, a clone was selected, and the insert of its plasmid was sequenced for confirmation. pUCSE isolated from the clone was then digested with Eco RI and Hind III using the manufacturer's protocol and the large fragment was isolated. The following adaptor (SEQ ID NO: 14 and 37, respectively) was ligated to the large fragment to give plasmid pUCSE-D1 which contained the first di-word (underlined).

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          Bse RI
EcoRI PstI      BbsI      Bsp120I      HindIII
  ↓    ↓    ↓          ↓          ↓          ↓
aattctgcagaggagatgaagacgaaaagaaaggggcccattgctgca
      gacgtctcctctacttctgcttttctttccccgggtacgacgttcga
                                   ↑
                                   BbvI

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Formula I

Further plasmids, pUCSE-D2 through pUCSE-D64, containing di-words were separately constructed from pUCSE-D1 by digesting it with Pst I and Bsp 120 I and separately ligating the following adaptors (SEQ ID NO NOs: 15 and 39, respectively) to the large fragment.

gaggagatgaagacga [word] [word] g
acgtctcctctacttctgct [word] [word] cccgg

Formula II

The words of the top strand were selected from the following minimally cross-hybridizing set:
.gatt, tgat, taga, ttg, gtaa, agta, atgt, and aaag. After cloning and isolation, the inserts of the vectors
were sequenced to confirm the identities of the di-words.